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Identification, Sequencing, Expression and Evolutionary Relationships of the 11S Seed Storage Protein Gene in *Chenopodium quinoa* Willd.

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IDENTIFICATION, SEQUENCING, EXPRESSION AND EVOLUTIONARY
RELATIONSHIPS OF THE 11S SEED STORAGE PROTEIN GENE IN
CHENOPODIUM QUINOA WILLD.

by

Marie Renee Barrett Balzotti

A thesis submitted to the faculty of

Brigham Young University

in partial fulfillment of the requirements for the degree of

Master of Science

Department of Plant and Animal Sciences

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BRIGHAM YOUNG UNIVERSITY

GRADUATE COMMITTEE APPROVAL

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ABSTRACT

IDENTIFICATION, SEQUENCING, EXPRESSION AND EVOLUTIONARY RELATIONSHIPS OF THE 11S SEED STORAGE PROTEIN GENE IN *CHENOPODIUM QUINOA* WILLD.

Marie R.B. Balzotti

Department of Plant and Animal Sciences

Master of Science

Quinoa (*Chenopodium quinoa*) is an Andean crop adapted to harsh environmental conditions and containing a high and well-balanced protein composition. Two seed storage proteins, the 2S albumin and 11S globulin, are the major amino acid reservoirs for the developing seedling. An in-depth study of the genes encoding these proteins is necessary to understand the source of quinoa's protein quality. Our objectives include identification and sequencing of the 11S genomic and cDNA clones, analysis of 11S expression profiles in different quinoa accessions and evaluation of evolutionary relationships between the sequence of the 11S gene in quinoa and homologous genes in other species.

Clones containing 11S cDNA and genomic DNA were identified and sequenced. Two copies of the gene were found to be present at two different loci in the quinoa genome. The amino acid composition of the 11S gene was also analyzed. Results show that the 11S gene contains a well-balanced assortment of essential amino acids with

relatively high levels of glutamate/glutamine, aspartate/asparagine, arginine, serine, leucine and glycine, typical of other 11S seed storage proteins.

Total RNA and globulin was extracted from seed collected at different developmental intervals from nine quinoa accessions. Expression profiles were determined by evaluating 11S transcript levels using relative quantification real time RT-PCR and comparing relative 11S globulin accumulation using sodium SDS-PAGE. The 11S gene was found to be expressed during late-maturation regardless of differences in maturation rate.

A portion of the amino acid sequence of the 11S and homologous genes was chosen for phylogenetic analysis. Fifty five such sequences from 50 different plant species were assembled and aligned. Two phylogenetic reconstructions, one using the parsimony method and the other using Bayesian analysis, were generated in order to analyze evolutionary relationships between the 11S gene in quinoa and homologous genes in other species. Relationships shown by both reconstructions for sequences from closely-related species were consistent with taxonomic clustering. Both reconstructions showed less resolution involving relationships between distantly-related angiosperm taxa indicating a wide divergence of sequence at the angiosperm level and a need for additional angiosperm sequence data for finer resolution.

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Isolation of BAC clones containing the 11S gene from a BAC library for *Chenopodium quinoa* Willd.

ABSTRACT

Quinoa is a crop from the Andean region of South America. Its seed has an unusually well-balanced amino acid composition and the plant is well-adapted to harsh environmental conditions including frost, drought and saline soils. The objective of this study was to identify BAC clones containing the 11S seed storage protein, one of two major storage proteins found in quinoa seed using a quinoa BAC library. Seven positive clones were identified from the library using an 11S cDNA probe. Southern blotting was performed using BAC DNA and genomic DNA from four accessions, all digested with *EcoRI*. Two copies of the 11S gene were identified which represent two distinct loci within the quinoa genome.

INTRODUCTION

Quinoa (*Chenopodium quinoa*) is a putative allotetraploid ($2n = 4x = 36$) and a member of the Amaranthaceae (or Chenopodiaceae) family. Other members of this family having some commercial importance are amaranth (*Amaranthus* spp.), spinach (*Spinacea oleracea* L.) and sugar beet (*Beta vulgaris* L.). Quinoa inhabits a variety of environments from the high Andean region to valleys and coastal areas and was an important crop of the Incan empire (Galwey, 1995). After the Spanish conquest, quinoa cultivation was discouraged due to its affiliation with cultural and religious practices and

replaced by introduced species such as wheat (*Triticum aestivum* L.) and barley (*Hordeum vulgare* L.). Quinoa cultivation was preserved throughout the years only by poor or isolated indigenous communities and small farms where it was able to maintain high genetic diversity (Castillo, 1995). Recently, there has been a revival of quinoa production and renewed interest has spread outside the Andean region. Commercially grown quinoa seed has been successfully promoted not only in South America but also in Europe, Australia, North America and Japan. Quinoa's ability to adapt to drought, frost and alkaline or salty soil conditions as well as its high protein content and favorable assortment of amino acids has made it an attractive candidate for crop improvement and attracted interest in the fields of genetic and molecular research.

There have been several studies involved in the characterization of quinoa's seed storage proteins. Quinoa seed proteins were electrophoretically characterized in order to create genetic markers for identification and classification of quinoa accessions as well as a resource for phylogenetic and genetic variability studies (Fairbanks et al., 1990). An analysis of quinoa seed fractions including protein analysis was conducted by Chauhan et al. (1992). Brinegar and Goundan (1993) specifically characterized individual seed storage proteins with the isolation and characterization of the 11S seed storage protein, which they call chenopodin. In their study, protein structure, amino acid compositions, and the N-terminal sequence for the basic subunit were determined. In addition, the 2S seed storage albumin, the other major protein component of quinoa, was isolated and characterized (Brinegar et al., 1996).

Since the discovery of quinoa's specific seed storage proteins, recent studies have provided genetic tools for further characterization of the genes encoding these proteins.

Over 400 simple sequence repeats were characterized by Mason et al. (2004) in order to provide groundwork for the development of a genetic linkage map containing 230 amplified length polymorphism (AFLP), 19 simple sequence repeat (SSR), and 6 randomly amplified polymorphic DNA (RAPD) markers (Maughan et al., 2004). Coles et al. (2004) reported the construction of a quinoa cDNA library from seed and floral tissues and its utility in the identification of 51 single nucleotide polymorphisms (SNPs) and 424 expressed sequence tags (ESTs) from which the clones containing 11S cDNA were identified and sequenced. And finally, a BAC library for quinoa was developed and reported by Stevens et al. (2006) containing 74,496 clones with approximately 9x coverage of the quinoa genome.

One utility of the BAC library is sequence detection and isolation. Here we report the identification of clones containing putative 11S seed storage protein genomic DNA from the quinoa BAC library.

RESULTS

Identification and characterization of seed storage proteins in quinoa was an important first step in understanding how protein is stored in the seed in such high quantities and with such a well-balanced assortment of essential amino acids. In order to identify clones of the 11S gene, double spotted high-density blots representing the quinoa BAC library were hybridized with a radiolabeled fragment of the 11S gene. Ten clones hybridized with the 11S probe and seven of these were identified and selected from the BAC library.

Figure 1 shows the results of Southern blotting of *EcoRI*-digested BAC and genomic DNA probed with a fragment of the 11S gene. The radiolabeled probe was developed using primers specific to the 11S cDNA clone and amplifying a portion of it from BAC DNA extracted from one of the positive clones, 77L9. Lanes 1-7 include the seven BAC clones: 77L9, 13H24, 36L11, 82M10, 119J6, 155I19, and 164F2, respectively. Lanes 8-11 include genomic DNA from four homozygous lines: Chucapaca, KU-2, NL-6, and 0654, respectively. These homozygous are parentals of populations used for genetic mapping (Maughan et al., 2004). Two different bands, 4 and 4.5kb, were observed in lanes representing the seven BAC clones, similar in size to the bands of Chucapaca and 0654, two highland ecotypes. In addition, Southern blotting with 'Real', a highland variety and the quinoa accession used to construct the BAC library (Stevens et al., 2006), showed similar banding to Chucapaca and 0654 (data not shown). The 4.5kb band appears to be monomorphic across the four accessions whereas the 4kb band in the highland varieties and the 5.5kb band in the lowland varieties indicate a polymorphism in the *EcoRI* restriction site between the highland and lowland ecotypes.

DISCUSSION

Following construction of a BAC library, clones containing the 11S gene were identified and characterized. Restriction enzyme digestion followed by Southern blotting revealed two distinct bands evident in four parental accessions and a single band either 4kb or 4.5kb in size in the BAC clones. The relatively small size of the fragments suggests that the 11S gene may be present as a single copy at two separate loci within the genome. This notion is further supported by the fact that the two copies of the gene show

up in separate BAC clones and are located within separate linkage groups on the developing quinoa map (Jarvis, 2006). The restriction fragment length polymorphism identified between the highland and lowland ecotypes will be useful for genetic marker development. In addition, because quinoa is a putative allotetraploid and thought to have arisen from two diploid ancestors, it is possible that future mapping studies may show these two genes to be on homeologous chromosomes providing valuable information for studies involving quinoa ancestry. Quinoa's extraordinarily nutritional quality and ability to tolerate a wide variety of environmental conditions make it a valuable commodity. Understanding the proteins responsible for quinoa's unique amino acid composition is essential to future improvements in the quantity and quality of its seed storage proteins.

MATERIALS AND METHODS

Plant materials

Tissue for use in DNA extractions were collected from the first two true leaves of two-to-four-week-old plants grown under greenhouse conditions at Brigham Young University (BYU) of four inbred lines: NL-6, KU-2, 0654, and Chucapaca. Harvested tissue was immediately frozen and stored at -80°C.

Hybridization of the quinoa BAC library

An 11S gene probe was developed by amplification of a fragment of the gene with the following primers: forward 5'- GCACTAGTGTTGTTGAATGGGTGC -3' and reverse 5'- GCGTTCACCCCTCTGGGATT -3' derived from the cDNA sequences (GenBank accession AY562550). Hybridization of the BAC library was carried out according to the method described by Stevens et al. (2006).

DNA extraction and Southern blotting

Positive clones from the BAC library screening were grown in LB media containing 12.5 µg/ml chloramphenicol and BAC DNA was extracted using a BACMAX™ DNA purification kit (Epicentre, Madison, WI). Genomic DNA was extracted from NL-6, KU-2, 0654, and Chucapaca according to the protocol reported by Saghai-Marroof et al. (1984). Genomic and BAC DNA was digested using *EcoRI* and fragments were separated on a 0.8% agarose gel by electrophoresis. The gel was blotted using downward capillary transfer to a Hybond™-N+ positively charged nylon membrane (Amersham, Piscataway, NJ) and hybridized with a ³²P-labeled 11S probe at 60°C overnight. The membrane was washed and visualized the same as described by Stevens et al. (2006).

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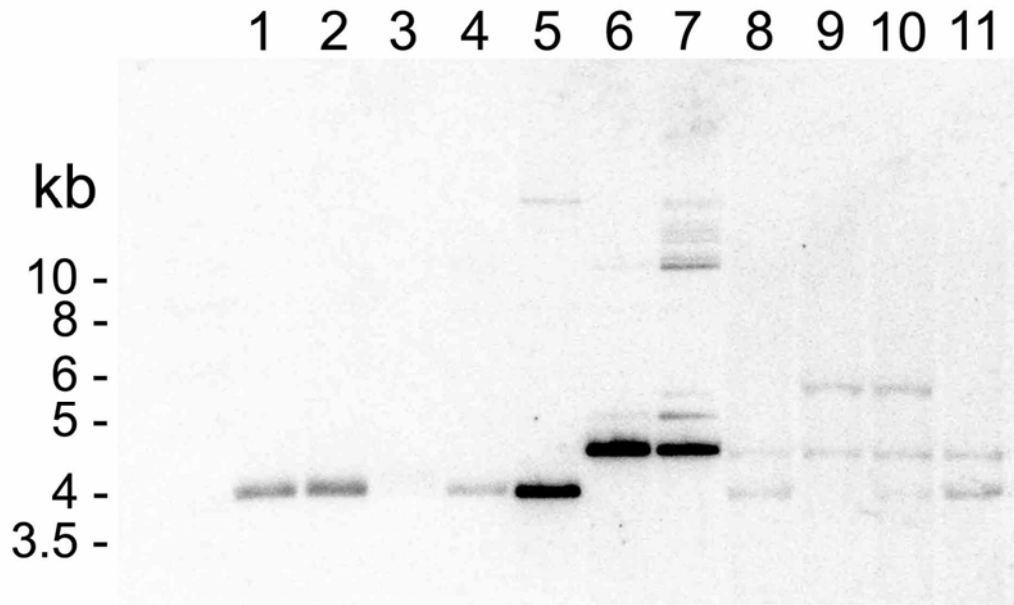


Figure 1. Southern blot of quinoa BAC DNA containing the 11S gene and genomic DNA from four quinoa accessions from inbred lines; lanes 1-7: 36L11, 82M10, 119J6, 155I19, and 164F2, respectively; lanes 8-11: genomic DNA from Chucapaca, KU-2, NL-6, and 0654 respectively. All DNA was digested with *EcoRI* and probed with a fragment of the 11S seed storage protein gene. The sizes of DNA markers are given in kilobases (kb) shown on the left. Lane 3 containing 119J6 BAC DNA shows a faint band at 4kb due to poor DNA concentration. Additional bands are shown in lanes 5-7 which is most likely due to incomplete digestion.

Sequencing, expression and evolutionary relationships of the 11S seed storage protein gene in *Chenopodium quinoa* Willd.

ABSTRACT

Quinoa (*Chenopodium quinoa* Willd.) is a pseudocereal that provides nutritional value for high Andean plateau farmers and commercial value for many South American countries. Present interest in quinoa is due to its tolerance of harsh environments, its well-balanced amino acid composition and nutritional value. The seed storage proteins of quinoa, particularly the 11S globulins and 2S albumins are responsible for providing a reservoir of nitrogen and amino acids for the developing seedling upon germination. The genomic and cDNA sequences for two copies of the 11S gene at two different loci were determined. Transcription initiation and termination sites, coding regions, and introns were determined by comparing genomic to cDNA sequences. Gene expression and protein accumulation data were obtained using real time RT-PCR and separation by SDS-PAGE, respectively. Using the coding DNA sequence for the well-conserved 11S basic subunit, phylogenetic relationships were analyzed between quinoa and other species for which 11S and homologous sequences have been reported.

INTRODUCTION

Quinoa (*Chenopodium quinoa* Willd.) is an allotetraploid dicotyledonous pseudocereal whose seed is used as a source of nutrition for people in the Andean region to which it is indigenous. It has been an important cultural and dietary component for people of the Andean region since 3000 B.C. and continues to be a major resource for the Quechua and Aymara people (Brinegar, 1997). Quinoa is able to tolerate a wide variety of environments from valleys and coastal foothills to the semi-arid and cool Altiplano. In addition, approximately 12.6-13.7% of the dry weight of quinoa seed is protein (Cardoza and Tapia, 1979; Chauhan et al., 1992) which is high compared to most important cereal crops such as rice, corn and millet (Charalampopoulos, 2002). In addition, its amino acid composition exceeds the FAO recommendations for human protein consumption (Ruas et al. 1999) with relatively high quantities of lysine and sulfur-containing amino acids which are usually limiting in the seeds of cereals and legumes, respectively. Seed storage proteins play a vital role in the protein composition of the seed. It is therefore important to understand the genes encoding these proteins, their expression profiles and how they relate to analogous genes in other species in order to understand how protein is deposited in the seed and why quinoa, specifically, contains a relatively high quantity and well balanced assortment of amino acids.

Large quantities of seed storage proteins accumulate in developing seeds, as a source of nitrogen, carbon, sulfur and amino acids for use in germination and growth of the developing seedling. These protein reserves are stored in the cells of the endosperm and embryo in protein storage vacuoles or specialized aggregates called protein bodies, assembled within the endoplasmic reticulum (Prego et al., 1998). Seed storage proteins

are generally classified according to their solubilities: albumins, globulins, prolamins, and glutelins which are soluble in water, saline solutions, alcohol, and alkali solutions respectively (Osborne, 1924). Storage proteins are further classified based on their sedimentation coefficients ($S_{20,w}$). Globulins generally fall into two major groups based on these coefficients: the 7-8S vicilin-type and the 11-12S legumin-type. Since legumin-type seed storage proteins vary in size, we will refer to the 11-12S globulins collectively in other species as legumins. The 11S globulin, therefore, is a sub-type of the legumin seed storage proteins.

Recently, efforts have been made to understand the legumin gene families and the structures of the proteins they encode. Legumin sequences have been reported for economically important seed crops such as rice (Okita et al., 1989; Wen et al, 1989; Takaiwa et al., 1991), oat (Shotwell et al, 1988; Schubert et al., 1990; Tanchak et al., 1995), maize (Woo et al., 2001; Yamagata et al., 2003), and soybean (Momma et al., 1985; Utsumi et al., 1987; Nielsen et al, 1989; Xue et al., 1992). The crystal structure of an 11S globulin from soybean was determined, providing new insights into conserved binding domains, quaternary structure and how structure relates to packaging, storage and degradation (Adachi et al., 2003).

The 11S globulin is a hexameric protein consisting of six pairs of acidic and basic subunits with each subunit pair connected by a disulfide bond. It is translated as a single precursor polypeptide containing acidic and basic subunits as well as a signal peptide responsible for translocation of the precursor into the endoplasmic reticulum (ER). This signal peptide is cleaved upon entry into the lumen of the ER (Herman and Larkins, 1999). The 11S protein assembles as a trimer in the ER which is transported to the

vacuoles through the Golgi apparatus. Further processing occurs at a well-conserved cleavage site by an asparaginyl endopeptidase which divides the subunit into its acidic and basic components and hexamer formation finally takes place. The assembly of the hexamer is a direct result of this second cleavage event (Dickinson et al., 1989). At the onset of germination, degradation results due to pH-influenced flexibility and dissociation of the 11S hexamer as well as the actions of vacuolar-targeted proteinases formed by germination-related regulatory mechanisms (Adachi et al., 2003; Shutov et al., 2003).

Conserved regions within the 11S amino acid sequence are necessary for proper folding, packaging, storage, and proteolysis. Temporally controlled protein protection and degradation is determined by specific structural features. Distance-based phylogenetic analysis of legumin-encoding sequences as well as a review of the major groups of legumins was reported by Häger and Fischer (1999). The origin and early evolution of legumin structures have been reported by Shutov and Baumlein (1999) who describe their evolution from a germin-like ancestor. Following divergence of seed storage proteins into vicilin and legumin classes, each acquired independently their storage-related properties. Seed storage and analogous proteins seem to have an ancient history, present in the early progenitors of plants including mosses and fungi (Shutov et al., 2003). Thus, these gene sequences may indeed prove useful as phylogenetic markers in molecular evolution studies as well as provide insight into evolutionary modifications under the selective constraints of either better storage capabilities or degradation. In addition, the study of conserved and variable regions and how they relate to structure and

metabolism may provide valuable information into how these protein sequences may be altered and improved for nutritional purposes in quinoa and other species.

The 11S globulin and the 2S albumin are the two major storage proteins in quinoa seed (Brinegar, 1997). The 11S globulin in quinoa has a typical 11S quaternary structure with an estimated native molecular weight of 320 kDa. The acidic and basic polypeptides have average relative molecular weights of 35,500 Da and 22,500 Da, respectively (Brinegar and Goundan, 1993).

Here we report the genomic and cDNA sequences from the quinoa 11S gene. We present data on 11S gene expression during seed development and the corresponding accumulation of 11S globulin within the seed. Coding DNA sequences were used to conduct a phylogenetic study between the 11S sequence in quinoa and homologous seed storage proteins of diverse species. The results of this analysis may answer questions concerning legumin molecular evolution, why quinoa contains such an extraordinary protein content compared to important cereal crops and how the 11S sequence of quinoa is different from or similar to those of other species.

RESULTS

Isolation and Characterization of the Quinoa 11S Gene

The cDNA sequences for 17B7 (figure 1) and 8B14 (figure 2) were determined by primer walking off clones reported by Coles et al. (2004). Isolation of BAC clone 77L9 was previously reported by Stevens et al. (2006) and the same protocol was used in the recent identification of BAC clone 164F2. These two clones contain putative 11S genes from two separate loci within the quinoa genome. The nucleotide sequence of the entire

gene and flanking regions were obtained directly from the BAC clones using a primer walking strategy. The nucleotide sequence of the 11S gene region from clones 77L9 and 164F2 are shown in figures 3 and 4, respectively. Coding and flanking regions of the genomic nucleotide sequences of 77L9 and 164F2 had 99.5% and 99.8% sequence identity with that of the cDNA sequences of 17B7 and 8B14, respectively.

Initiation and termination codons and intron positions were determined by comparing genomic and cDNA sequences. Three introns were identified within the 11S gene and are shown in Figs. 3 and 4. The locus corresponding to BAC clone 164F2 and cDNA clone 8B14 contain a 1437 base pair open-reading frame that encodes a polypeptide 479 amino acids in length, whereas the locus corresponding to BAC clone 77L9 and cDNA clone 17B7 contain a 1440 base pair reading frame coding for a polypeptide with 480 amino acids. The three-base difference between the two clones results in an arginine insertion at position 285. Although the coding and flanking regions of genomic nucleotide sequences 77L9 and 164F2 share approximately 97% and 92% sequence identity, respectively, intron sequence identity between the two was only 57% on average.

The 11S amino acid sequence in quinoa is similar to 11S seed storage proteins found in other species. Amino acid compositions of polypeptides from the two loci are shown in Table 1. The first 25 hydrophobic amino acid residues are typical of a signal peptide found in other legumin-like seed storage proteins that is removed once entering the lumen of the ER (Utsumi, 1992). An additional processing site (between asparagine 293 and glycine 294 in 17B7 and asparagine 291 and glycine 292 in 8B14) in which the precursor polypeptide is separated into acidic and basic subunits is shown in figures 1-4. This

cleavage event would result in acidic and basic subunits 267-268 and 188 amino acids in length respectively. Other conserved sequences include regions surrounding amino acid residues that participate in covalent and hydrogen binding. Four of these regions contain cysteine residues which participate in disulfide bond formation in other species (Adachi et al., 2003).

11S Gene Expression and Protein Accumulation

To determine the 11S gene expression profile in quinoa, seeds were collected at eight- and ten-day intervals after anthesis until dessication from nine quinoa accessions: NL-6, KU-2, Mocko, Maniqueña, Ollague, Ratuqui, Sayaña, Chucapaca, and 0654. Accessions were chosen based on geographic location and maturation rate. Plants were selected from coastal regions and the Altiplano (high Andean plateaus) and are shown in Table 2. Altiplano types are subdivided into Salares (areas containing salt flats) and Valley. Maturity was determined at the time 90% of the seeds were fully dessicated.

Following seed collection, total RNA was extracted and quantified. Relative quantification of 11S mRNA in seeds from different developmental stages was determined by one-step reverse transcriptase real time PCR using the GAPDH gene as an endogenous control. In order to confirm expression at the protein level, relative 11S globulin quantity was determined by SDS-PAGE analysis. Figure 5 shows the results of 11S mRNA and globulin quantification at specific days after anthesis (DAA). Few 11S mRNA transcripts were detected during early seed development in all accessions. In general, 11S expression was detected followed by a substantial accumulation of 11S protein. As seeds became mature, expression decreased and protein accumulation remained high.

Based on our observations of plants grown under the same greenhouse conditions, five types of maturation rates were identified according to days to desiccation, 11S RNA and protein accumulation patterns and are listed together with accession names and locations in Table 2. Type I consists of accessions KU-2 and NL-6 from coastal regions showing peak gene expression and 11S globulin accumulation at 24 DAA and maturity between 32 and 40 DAA. Type II, including Maniqueña and Mocko, two varieties located in the Salares, had peak gene expression at 24 DAA, detectable 11S globulin quantity occurring 32 DAA and maturity achieved between 40 and 50 DAA. Ollague (also located in the Salares), Ratuqui and Sayana (two accessions found in the Altiplano) were included as type III accessions and showed peak 11S transcript levels and substantial 11S globulin accumulation at 40 DAA and reached maturity between 50 and 60 DAA. Chucapaca, also located in the Altiplano, and 0654 from valleys, were included in type IV and type V, respectively, showed detectable 11S protein quantities at 50 DAA. However, Chucapaca reached maturity at approximately 60 DAA and 11S transcript levels reached a maximum amount 40 DAA while those of 0654 did so at 50 DAA and reached maturity between 60 and 70 DAA. As shown in Figure 5, the increases in 11S globulin levels in these different varieties appear to be correlated with the expression profiles for the 11S gene.

Phylogenetic Analysis of 11S Seed Storage Proteins

The coding DNA sequences of complete legumin sequences from 50 different plant species were translated, aligned and evaluated for phylogenetic signal quality. Common name, species, families, and orders for the plant species utilized in the phylogenetic analysis are presented in Table 3. Alignments were created using whole or portions of

the legumin sequence. Using distance-based and maximum parsimony methods for tree reconstruction in MEGA3 (Kumar et al., 2004) preliminary trees were evaluated to assess the quality of the alignments. Amino acid alignments generated from only the basic subunit sequence of the legumin gene (partial alignment shown in Figure 6) resulted in detectable amino acid conservation, consistency between trees generated by the aforementioned methods and taxonomic clustering. It was therefore selected for further analysis.

Figures 7 and 8 show parsimony and Bayesian tree reconstructions, respectively. In both reconstructions, with posterior probabilities and bootstrap values of 100, the two quinoa 11S sequences formed a monophyletic group with that of a *Amaranthus hypochondriacus* sequence (Genbank accession X82121), the only other available sequence from a species within the Amaranthaceae family. Using amino acid alignments so identities and positives were maximized, the 11S basic subunit showed over 74% sequence identity between amaranth and quinoa. Other monophyletic and well-resolved taxonomic groups represented in both trees include taxa from the Poaceae, Brassicaceae, Cupressaceae, and Pinaceae families, the Fagales order and the clade for gymnosperms. As expected, the sequences from members of the Fabaceae family formed two monophyletic groups, each containing 11S sequences derived from different subfamilies that diverged early in angiosperm evolution (Yagasaki, 1997). The phylogenetic relationships between the gymnosperm outgroup taxa are well-resolved in both trees and reveal a clear bifurcation of angiosperm and gymnosperm legumin genes. Although both reconstructions show similar relationships between taxa, the Bayesian consensus tree has better support, indicated by high posterior probability values, than the parsimony tree for

unresolved portions of the tree including relationships between taxa from species of different families and orders.

DISCUSSION

11S Gene Sequences

Two BAC clones, 77L9 and 164F2, containing 11S genomic DNA and two cDNA clones, 17B7 and 8B14, were sequenced and analyzed. The coding sequences for 77L9 and 164F2 highly correlated with 99.7% and 99.9% identity to 17B7 and 8B14, respectively. The quinoa cultivar ‘Real’ was used in construction of both cDNA and BAC libraries. However, it is not truebreeding. Therefore, the slight difference in DNA sequence between the BAC and cDNA sequences is likely due to heterogeneity in the ‘Real’ line. Although the coding and flanking regions were well conserved between the two loci (approximately 96% identity overall), the introns only correlated with approximately 57% identity.

The amino acid sequence of the 11S gene in quinoa is very similar to that of other species in domains necessary for binding and hexamer formation. Sequence similarities of the first 25 amino acid residues between quinoa and other species suggest that this region is a signal peptide that localizes the precursor to the ER where it is subsequently removed (Utsumi, 1992). Brinegar (1993) reported the N-terminal sequence of the basic subunit indicating that the polypeptide is indeed cleaved at a well-conserved site reported in other species (Utsumi, 1992). The sequence similarities of six binding regions suggest that the quinoa 11S hexamer has a structure similar to glycinin, for which the crystal structure has been determined (Adachi et al., 2003). Four of these binding regions

contain cysteine residues that participate in disulfide bond formation, one intrachain disulfide bond which occurs within the acidic polypeptide between the first two cysteine residues and one interchain disulfide bond which occurs between the last cysteine residue within the acidic subunit and the first cysteine in the basic subunit (Utsumi, 1992). (See Figures 1-4.)

The 11S amino acid sequence indicates a well-balanced composition and a high level of essential amino acids. Typical of 11S storage proteins, chenopodin contains high levels of glutamate/glutamine, aspartate/asparagine, arginine, serine, leucine and glycine (Brinegar, 1993). However, we note that the 11S gene is only one of two major seed storage proteins in the quinoa seed and that it is the combined contribution of the 11S and the 2S seed proteins that produce quinoa's unique seed protein quantity and composition. Thus, cloning and sequencing of the 2S albumin will be necessary to gain a complete understanding of the relative contributions of each storage protein within the developing seed and how quinoa seed accumulate such a high quantity of protein and essential amino acids.

Expression Patterns of 11S Genes in Different Quinoa Accessions

The gene expression and protein data presented here suggests that the accumulation of seed storage protein is associated with maturation rate. Early maturing varieties KU-2 and NL-6, for example had the first substantial accumulation of 11S mRNA (16 DAA) and protein (24 DAA) while late maturing variety 0654 did not show increased transcript levels and protein accumulation until 50 DAA. These data and those from the other quinoa accessions suggest that high expression of 11S mRNA occurs, on average, during late-maturation, consistent with the pattern of storage protein

accumulation during seed development and those of other 11S seed storage proteins (Nakamura, 2004). In this analysis, we have observed relative increases in 11S transcript levels and globulin. However, a quantitative study into the total protein content per seed of these and other quinoa accessions will be needed to determine if the timing of gene expression has an effect on the overall quantity of seed protein. Also, while analyzing plants grown in the greenhouse is important and informative, this study should be repeated in the field to see if our observations are consistent with those of field-grown plants.

Phylogenetic Relationships between Legumins of Various Species

The legumin coding DNA sequence of the basic subunit was used in order to analyze phylogenetic relationships between the 11S gene of quinoa and other species. The basic subunit of legumin is well-conserved and contains sequence elements that are necessary for its proper assembly and packaging. These include the first N-terminal amino acids involved in the recognition of the peptide cleavage site between the acidic and basic subunits followed by an amino acid sequence which forms a β -barrel structure. Also included are conserved regions surrounding several amino acids involved in hydrogen bonding, hydrogen-bonded salt bridges and a cysteine residue involved in disulfide bond formation (Adachi et al., 2003). Thus the entire legumin basic subunit sequence or portions of the basic subunit are suitable phylogenetic sequences for studies involving the molecular evolution of seed storage proteins (Fischer et al., 1996; Häger and Wind, 1997; Shutov and Bäumlein, 1999).

Two phylogenetic trees, one based on maximum parsimony and another on Bayesian analysis, are shown in figures 7 and 8, respectively. These two methods, which

utilize very different assumptions, were selected in order to validate the relationships shown in the reconstructions. Maximum parsimony assumes that the best tree is one in which evolutionary steps are minimized while Bayesian analysis implements a model of evolution in addition to the sequence data in order to maximize the probability of a tree. Because one method lacks outside parameters assuming that one evolutionary event is just as likely as another and the other uses established parameters in order to predict associations, we assumed that relationships established by both reconstructions would be well-supported. Indeed, all monophyletic groups defined by the maximum parsimony tree were also supported by the Bayesian tree.

The data suggest that the Bayesian analysis was better suited for resolving relationships between more distantly-related taxa than the parsimony method. Indeed, most unresolved portions of the Bayesian reconstruction angiosperm backbone had posterior probabilities just under 90 and are therefore not shown in Figure 8. A third and fourth node with posterior probabilities of 83 and 88, respectively was present following the bifurcation of lower angiosperm legumin sequences in ginger, mint and yam (Fischer et al., 1996). The third node represented the divergence in a sequence from arrowhead. The fourth node separated two clades from the remaining angiosperms, one including sequences from members of the Poaceae family, magnolia, african oil palm, almond and coffee and the other composed of sequences from sesame, pumpkin, orange and buckwheat (data not shown). Although these relationships are based on posterior probabilities under 90, they are still supported by the majority of the data.

Following the divergence between gymnosperms and angiosperms, there appears to be an explosion of sequence divergence in the legumin genes as evident by the

presence of multigene families and the lack of resolution in the phylogenetic tree reconstruction. Multigene families for the legumin-like genes have been identified in many species including those of legumes (Domeney et al., 1986; Heim et al., 1994; Nielsen et al., 1989), brassicas (Breen and Crouch, 1992; Depigny-This et al., 1992; Pang et al., 1988) and cereals (Okita et al., 1989; Shotwell et al., 1988). Multigene families have also been identified in gymnosperms (Hager et al., 1995; Wind and Hager, 1996; Hager and Wind, 1997). However, these sequences have a much higher degree of homology with each other than those of angiosperms. There does not appear to be an 11S multigene family in quinoa (Stevens et al., 2006), nor have there been any identified in amaranth (Barba de la Rosa et al., 1996). However, other 11S genes or pseudogenes may be present in the quinoa genome but have not yet been identified due sequence dissimilarity. If additional 11S genes are present but nonfunctional, it would be interesting to understand what type of selection may have resulted in such an inactivation.

Both phylogenetic trees were well-resolved between closely-related taxa. Quinoa's 11S sequence placement with amaranth exhibited posterior probabilities and bootstrap values of 100. The observation that the quinoa sequence was located where it would be expected to fall (with the only other sequence from a member of the Amaranthaceae family) and the high degree of homology between it and amaranth suggests that selection on this sequence is absent or subtle or that opposing selective forces do not allow the sequence to change substantially. Indeed, Shutov et al. (2003) suggested that seed storage protein structure evolution is under the constraints of two antagonistic selective forces: selection for better storage capabilities and selection for improved degradation. Quinoa is a putative allotetraploid and we postulate that the two

copies of the 11S gene may be located on homeologous chromosomes and represent genes present in the diploid ancestors of quinoa (Stevens et al., 2006). Although the sequences for both loci are similar, transcript levels from each may not be equal. As an alternative, it is possible that the unusually high protein content of quinoa is a direct result of tetraploidization and the unaltered expression of both genes since polyploidy generally increases gene expression levels overall (Osborn et al., 2003). Thus, seeds containing higher protein reserves for germination and the developing seedling may have been at a selective advantage.

The ability to resolve relationships between sequences from gymnosperms and other ancient taxa is shown by both parsimony and Bayesian trees. We note, however, that the radiation at the angiosperm level as well as the lack in sufficient sequence data to match the broad range of angiosperm species creates some challenges when using the legumin sequence as a phylogenetic tool. Indeed, without adequate sequence data and the added complication of multigene families that may have arisen at different occasions during evolutionary time, it is difficult to know if homologous genes are being compared and taxa may be grouped inappropriately. However, it is evident from this study that there is sufficient phylogenetic signal even from diverse species to conduct future studies into molecular evolution of legumin genes in angiosperms as new sequences are discovered.

MATERIAL AND METHODS

Plant materials

Quinoa accession Real was grown under greenhouse conditions at Brigham Young University. Tissue for use in DNA extractions were collected from the first two true leaves of two-to-four-week-old plants. Harvested tissue was immediately frozen and stored at -80°C.

Nine quinoa accessions were planted simultaneously and grown under greenhouse conditions: NL-6, KU-2, Maniqueña, Mocko, Ollague, Ratuqui, Sayaña, Chucapaca, and 0654. Seed was collected at eight day intervals after anthesis until maturity from each accession and immediately frozen and stored at -80°C.

cDNA and genomic sequencing

Genomic DNA was extracted from quinoa accession Real according to the protocol reported by Saghai-Marooof et al. (1984). An initial fragment of the quinoa 11S gene was obtained by amplification of genomic DNA using degenerate PCR primers designed from a partial quinoa 11S amino acid sequence (Brinegar and Goundan, 1993) as well as an alignment of a cDNA sequence from *Amaranthus hypochondriacus* (Barba de la Rosa et al., 1996). The forward primer sequence was 5'-AATGGKGTGGARGARACYATTTGC -3' and the reverse 5'-TGTKKGC GTTKAGGTTSYAGTG -3'. This gene fragment was used as a probe to screen the quinoa developing seed cDNA library reported by Coles et al. (2004). Positive clones were selected and their identities confirmed by Southern blotting. Clones 17B7 and 8B14 were selected for sequencing. Two BAC clones containing 11S genomic DNA, 77L9 previously isolated by Stevens et al. (2006) and 164F2 subsequently

identified and representing a second locus were identified by screening of a quinoa BAC library. The 11S genomic inserts within clones 77L9 and 164F2 and the cDNA inserts within clones 17B7 and 8B14 were sequenced by primer walking at the Brigham Young University DNA Sequencing Center (Provo, UT).

RNA extraction and relative quantification

Nine quinoa accessions were germinated and grown under the same greenhouse conditions. Temperatures were 20-25°C during the day and 15-18°C at night and plants were exposed to 12h days. Seed collected at eight- and ten-day intervals after anthesis from these accessions were ground to a fine powder in liquid nitrogen. Total RNA was extracted using an RNeasy® Plant Mini Kit (Qiagen, Valencia, CA) and quantified using a NanoDrop® ND-1000 Spectrophotometer (NanoDrop Technologies, Inc., Wilmington, DE). Primers and probes were designed for the 11S gene and the GAPDH gene: 11S forward 5'- GGCGGTCGCTTCCAAGA -3', reverse 5'- TTGCGAAAATGTGGCCTTGAC -3', probe 5'- CCAACACCAGAAGATCA -3'; GAPDH forward 5'- GGTTACAGTCATTCAGACACCATCA -3', reverse 5'- AACAAAGGGAGCCAAGCAGTT -3', probe 5'- CGCTTCCTGTACCAC -3'. Using the GAPDH gene as an endogenous control, multiplexed RNA samples were quantified using a TaqMan® One-Step RT-PCR Master Mix Reagents Kit (Applied Biosystems, Foster City, CA) on a 7300 Real Time PCR System (Applied Biosystems, Foster City, CA). Each sample contained 0.5µg of total RNA. Thermocycler conditions included the following: samples were held at 48°C for 30min, heated to 95° and held for 10min, followed by forty repetitions of temperatures at 95°C for 15s and 60°C for 1min.

Globulin extraction, quantification and separation by SDS-PAGE

Fifty milligrams of seed collected at eight day intervals after anthesis from the eight quinoa accessions were ground to a fine powder and 500µl of water was added to each sample. The samples were shaken at 4°C for over an hour, centrifuged at 13,000xg, and the supernatant containing the albumin fraction removed. The pellet was washed twice using the same protocol and dried. To each sample, 400µl 0.5M NaCl/50mM Tris-HCl, pH 8.0 was added. The samples were shaken at 4°C for over an hour, centrifuged at 13,000xg and the supernatant containing the globulin fraction was collected into separate tubes. Globulin was quantified using a BCA™ Protein Assay Kit (Pierce, Rockford, IL) on a NanoDrop® ND-1000 Spectrophotometer (NanoDrop Technologies, Inc., Wilmington, DE). Globulin samples were diluted 1:2 with Laemmli Sample Buffer (BioRad, Hercules, CA) and separated with SDS-PAGE on Ready Gels (Biorad, Hercules, CA). Gels were fixed overnight according to the manufacturer's instructions and visualized using Flamingo™ Fluorescent Gel Stain (Biorad, Hercules, CA) under UV light.

Taxonomic sampling and sequence alignment

Legumin sequences from 50 different species of plants were assembled using MEGA3 software (Kumar et al., 2004). These species and their accession numbers are listed in Table 3. Coding DNA sequences were downloaded from GenBank. Using the amino acid sequence of the 11S seed storage protein from *Chenopodium quinoa* (GenBank accession AAS67037) as a query sequence, a BLASTP search was performed (Altschul et al., 1990). Only legumin sequences from different plant species were chosen with E-values less than 1e-10 as to include sequences substantially similar and likely

related by descent to the 11S gene in quinoa. Incomplete sequences, less than 75% of the average length of the legumin gene, and redundant sequences were eliminated because the C-terminal basic amino acid sequence was used for analysis. One coding DNA legumin sequence from each species, except for quinoa and species from the Fabaceae family was selected based on its similarity to the 11S sequence from quinoa. Due to a greater amount of sequence data available for members of the Fabaceae family, legumin sequences from two major gene types (Yagasaki, 1997) were selected if available. The two quinoa sequences included in the analysis corresponded to the coding DNA sequences of cDNA clones 17B7 and 8B14 (GenBank accessions AY562549 and AY562550, respectively). The legumin basic subunit, a well-conserved sequence region, was chosen to improve the alignment because distantly related sequences were being compared. The selected coding DNA legumin sequences were translated into amino acid sequences using MEGA3. Amino acid sequences were aligned with ClustalW using the BLOSUM matrix and used as a template for the nucleotide alignment. ClustalW performs multiple sequence alignments of nucleotide or protein sequences using a progressive alignment algorithm including a gap cost model which penalizes insertions and deletions according to a linear function and sequence similarity for construction of a guide tree (Thompson et al., 1994).

Phylogenetic Reconstruction

Trees were constructed using distance, maximum parsimony and Bayesian methods. Branch support of parsimony and Bayesian reconstructions was tested using bootstrapping and posterior probabilities, respectively. ModelTest (Posada and Crandall,

1998) was utilized in order to select the most appropriate nucleotide substitution model for molecular evolution.

Distance, maximum parsimony and Bayesian analysis are three kinds of tree reconstruction methods reliant on different assumptions. Neighbor-joining is a distance method that expresses distance as a fraction of sites that differ between sequences. Maximum parsimony assumes that the tree generated by the fewest number of sequence changes between taxa is the most likely one. Bayesian analysis relies upon the concept of posterior probabilities, based on a postulated model of evolution that represents the probability of the relationship given the data. Trees are generated based on the model and the sequence alignment, sampled periodically and used to compute a consensus tree.

ModelTest (Posada and Crandall, 1998) is a program designed to assign the most statistically appropriate model of evolution to the alignment under evaluation. It employs two statistical approaches to model selection, the likelihood ratio test (LRT) and the Akaike information criterion (AIC). The LRT is generated by pairwise comparisons between the likelihood score of nested alternative hypotheses under a complex model. The AIC compares several models concurrently to determine the model that best fits the data while penalizing increasing numbers of parameters in the model (increasing parameters will always generate a better fit for the data but doesn't always represent the best model). The general time-reversible (GTR) model assumes that the probability of changing from one base into another is the same in one direction as it is in the reverse direction. Therefore, among four nucleotides there are six possible substitution rates. Additional parameters allow probabilities to be assigned to each site for designation to a specific rate category.

Measures of branch support for parsimony and Bayesian reconstructions can be tested by using bootstrapping and posterior probabilities, respectively. Bootstrap values are expressed as probabilities that taxa within a clade are always grouped within that clade. Posterior probabilities express the probability of relationships given the data and the model of evolution.

The phylogenetic signal of the data was explored using distance tree reconstruction in MEGA3. Parsimony reconstruction was accomplished using PAUP* version 4.0b10 (Swofford, 2001) and a bootstrap analysis with 1000 replicates was conducted with bootstrap values over 70 representing supported relationships. A parameter-rich GTR+I+G model to be implemented into the Bayesian analysis was selected from 56 alternative models with ModelTest as the most appropriate nucleotide substitution model for the aligned sequences. Bayesian consensus tree reconstruction was accomplished using MrBayes version 3.0b4 (Ronquist and Huelsenbeck, 2003) using default values as the starting parameters. Five million tree generations were conducted and trees were saved every 1000th generation. The consensus tree was generated with a burn-in at 20,000 tree generations and posterior probabilities above 90 represented supported relationships. Members of the Coniferales, Ginkgoales and Gnetales were assigned as a monophyletic outgroup based on putative sequence similarity to ancient taxa (Shutov et al., 1998)

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1	gattcgccacaaattcaatatcctcctcctcaaaaaaacaagcaaaaaaaaaaagtaa	60
61	aaaATGGCTAAGTCTACTACTACATTGTTTCTTCTTAGTTGTTCAATAGCACTAGTGTGG	120
1	M A K S T T T L F L L S C S I A L V L	19
121	TTGAATGGGTGCATGGGCCAAGGAAGGATGCGAGAGATGCAAGGCAATGAGTGCCAAATT	180
20	L N G C M G* Q G R M R E M Q G N E C§ Q I	19
181	GACCGGCTTACCGCCTCGAACCGACGAACCGGATTGAGCTGAGGGTGGATTGACTGAG	240
40	D R L T A L E P T N R I Q A E G G L T E	39
241	GTGTGGGACACCCAAGACCAGCAGTTCCAATGCTCCGGTGTGTCTGTTATTAGACGTACC	300
60	V W D T Q D Q Q F Q C§ S G V S V I R R T	59
301	ATTGAGCCTAATGGTCTTCTTTTGCCTTCTTTCCTACTAGTGGCCCTGAGCTCATTACATT	360
80	I E P N G L L L P S F T S G P E L I Y I	79
361	GAGCAAGGGAATGGAATAAGTGGGCTAATGATCCCGGGATGCCCGAGACATTTGAATCA	420
100	E Q G N G I S G L M I P G C‡ P E T F E S	99
421	ATGTACAAGAATCATGGAGAGAGGTATGAAGCGAGGAATGAGGGGCGGTCGCTTCCAA	480
120	M S Q E S W R E G M K R G M R G G R F Q	119
481	GACCAACACCAAGAAGATCAGGCACCTCCGTCGAAGGCCACATTTTCGCAATGCCGGCTGGA	540
140	D Q H Q K I R H L R Q G H I F A M P A G	139
541	GTTGCTCACTGGGCTTACAACACCGGAAATGAGCCTCTTGTGTGCTTATCCTCATTTGAC	600
160	V A H W A Y N T G N E P L V A V I L I D	159
601	ACCTCTAACCATGCTAACCAACTTGACAAGGATTACCCCAAGAGATTCTACCTAGCTGGT	660
180	T S N H A N Q L D K D Y P K R F Y L A G	179
661	AAACCCCAACAAGAGCACAGTCGCCACCAGCACAGAGGTGGAGAATCCCAGAGGGGTGAA	720
200	K P Q Q E H S R H Q H R G G E S Q R G E	199
721	CGCGGCACCGCGGCAACGTATTTCAGTGGGCTAGGCCACCAAGACCATAGCTCAATCCTTC	780
220	R G S G G N V F S G L G T K T I A Q S F	219
781	GGAGTTAGTGAGGACATAGCCGAGAAGCTCCAAGCCGAGCAAGACGAGAGAGGGCAACATA	840
240	G V S E D I A E K L Q A E Q D E R G N I	239
841	GTCTTAGTGAAGGGTCTTTCATGTGATCAAGCCCCAAGCAGCAGGTATATGATGAT	900
260	V L V Q E G L H V I K P P S S R S Y D D	259
901	GAGAGGGAACAACCGCCCATCGTAGCCCCAAGGTCTAACGGCTTGAAGAGACCATATGC	960
280	E R E Q R R H R S P R S <u>N* G L E E T I C‡</u>	279
961	TCGGTCGCTTAGTGAACAACATCGACGAACCCTCTAAGGCCGATTTTACTCCCGGAA	1020
300	<u>S A R L S E N I D E P S K A D V Y S P E</u>	299
1021	GCTGTAGACTTACTACCCTTAACAGCTTCAACCTCCCCATCCTCAGCAACCTCCGCTTG	1080
320	A G R L T T L N S F N L P I L S N L R L	319
1081	AGTCTGAGAAAGGGTGTCTTACAGGAACCGCATCATGGCACCACACTACAACCTAAAC	1140
340	S A E K G V L Y R N A I M A P H Y N L N	339
1141	GCCACAGCATAATCTACGGTGTACGAGGAAGAGACGTATCCAATCGTGAACGCACAA	1200
360	A H S I I Y G V R G R G R I Q I V N A Q	359
1201	GGAAACTCGGTGTTTCGACGACGAGCTAAGACAAGGACAACACTAGTGGTAGTCCACAGAAC	1260
380	G N S V F D D E L R Q G Q L V V V P Q N	379
1261	TTCGCAGTGGTGAAGCAAGCCGGTGAAGGAGGGTTCGAGTGGATCGCCTTCAAGACATGT	1320
400	F A V V K Q A G E E G F E W I A F K T C	399
1321	GAGAACGCTTGTTCCAAACCTAGCAGGCCGAACATCCGCCATCCGTGCCATGCCCCTTA	1380
420	E N A L F Q T L A G R T S A I R A M P L	419
1381	GAAGTGATCTCAAACATCTACCAGATCTCCCGTGAACAGGCGTACCGCCTCAAGTTAGC	1440
440	E V I S N I Y Q I S R E Q A Y R L K F S	439
1441	CGTCCGAAACACCTCTTCCGCCCGAGAACCAAGGGCGCCAAGGAGGGATTGGCT	1500
460	R S E T T L F R P E N Q G R Q R R D L A	459
1501	GCTtaagttggaaaaaacaacaatgtagatgcatggcataatgggtcacattatatatca	1560
480	A	480
1561	tgtaatgtgaaggatttttaggtgttttttaataataggatggataacaaaataa	1620
1621	aaggtagtcctttagaaggtcccttgttttgaaggtgtagctttagttgagaggattgga	1680
1681	gctgttatgtataaacttgtcaaaaaaaaaaaaaaaaaaaaaaaaaaaaaa	1728

Figure 1. The nucleotide and predicted amino acid sequence of cDNA clone 17B7. The * indicates putative cleavage sites that divide the precursor polypeptide into a 25-amino acid signal peptide, 267-residue acidic and 118-residue basic subunits. The ‡ and § indicate cysteine residues predicted to participate in intrachain and interchain disulfide bonds, respectively. The 9 underlined amino acids indicate a well-conserved processing site involved in peptidase recognition. Untranslated portions are indicated in lowercase.

1	gattcgccacaataaataatccttctccaaaaatacaagaaaaaaaaagagagtaaaaa	60
61	TGGCTAAGTCTACTACTACATTGTTTCTTCTTAGTTGTTCAATAGCACTAGTGTGTTGA	120
1	M A K S T T T L F L L S C S I A L V L L	20
121	ATGGGTGCATGGGCCAAGGAAGGATGCGAGAAATGCAAGGCAATGAGTGCCAAATCGACC	180
21	N G C M G* Q G R M R E M Q G N E C S Q I D	40
181	GGCTTACCGCCCTCGAACCGACGTATCGGATTCAGGCTGAGGGTGGGTTGACTGAGGTGT	240
41	R L T A L E P T Y R I Q A E G G L T E V	60
241	GGGACACCCAGGACCAGCAGTTCCAATGCCTCCGGTGTGTCTGTTATTAGGCGTACCATTG	300
61	W D T Q D Q Q F Q C S S G V S V I R R T I	80
301	AGCCTAATGGTCTTCTTTTGCCTTCTTCTACTAGTGGCCCTGAGCTCATTTACATTGAGC	360
81	E P N G L L L P S F T S G P E L I Y I E	100
361	AAGGGAATGGAATAAGTGGGCTGATGATCCCGGGTGCCCGGAGACATTTGAATCAATGT	420
101	Q N H G I S G L M I P G C † P E T F E S M	120
421	CACAAGAATCATGGAGAGAGGGTATGGAGCGAGGAATGAGGGCGGTGCGCTTCCAAGACC	480
121	S Q E S W R E G M E R G M R G G R F Q D	140
481	AACACCAGAAGATCAGGCACCTCCGTCAAGGCCACATTTTCGCAATGCCGGCCGGAGTTG	540
141	Q H Q K I R H L R Q G H I F A M P A G V	160
541	CTCACTGGGCTTACAACCTCCGAAATGAGCCACTTGTGCTGTTATCCTCATTGACACCT	600
161	A H W A Y N S G N E P L V A V I L I D T	180
601	CTAACCATGCTAACCAACTTGACAAGGATTACCCCAAGAGATTCTACCTAGCTGGTAAAC	660
181	S N H A N Q L D K D I P K R F Y L A G K	200
661	CCCAACAAGAGCACAGTCGCCACCATCACAGGGGTGGAGAATCCCAGAGGGGTGAACACG	720
201	P Q Q E H S R H H H R G G E S Q R G E H	220
721	GCAGCGACGGCAACGTATTTCAGTGGCCTAGACACCAAGTCCGTAGTTCAATCCTTCGGAG	780
221	G S D G N V F S G L D T K S V V Q S F G	240
781	TTAGTGAGGACATAGCGGAGAAGCTCCAAGCCAAGCAGGACGAGAGAGGCAACATAGTCT	840
241	V S E D I A E K L Q A K Q D E R G N I V	260
841	TAGTGCAAGAGGGTCTTCATGTGATCAAGCCCCAAGCAGCAGGTCCATATGATGATGAGA	900
261	L V Q E G L H V I K P P S S R S Y D D E	280
901	GGGAACAGCGCCATCGTAGCCCAAGGTCTAACGGCTTGGAAAGAGACCATATGCTCGGCTC	960
281	R E Q R H R S P R S N* G L E E T I C † S A	300
961	GCCTTAGTGAGAACATCGACGATCCCTCTAAGGCCGATGTTTACTCCCCGGAAGCTGGTA	1020
301	S L S E N I D D P S K A D V Y S P E A G	320
1021	GACTTACCACCCTTAACAGCTTCAACCTCCCATCCTCAGCAACCTCCGCTTGAGTGCTG	1080
321	R L T T L N S F N L P I L S N L R L S A	340
1081	AGAAGGGTGTCTCTACAGGAACGCAATCATGGCACCACACTACAACCTAAACGCCACACA	1140
341	E K G V L Y R N A I M A P H Y N L N A H	360
1141	GCATAATCTATGGTGTACGAGGACGAGGACGTATCCAAATCGTGAACGCACAAGGAAACT	1200
361	S I I Y G V R G R G R I Q I V N A Q G N	380
1201	CAGTGTGACGATGAGCTAAGACAAGGACAACCTAGTGGTAGTCCCACAGAACTTCGCAG	1260
381	S V F D D E L R Q G Q L V V V P Q N F A	400
1261	TGGTGAAGCAAGCCGGTGAGGAAGGATTCGAGTGGATCGCCTTCAAGACATGTGAGAACG	1320
401	V V K Q A G E E G F E W I A F K T C E N	420
1321	CTTTGTCCAAACCCTAGCAGGCCGAACATCCGCCATACGTGCCATGCCCGTAGAAGTAA	1380
421	A L F Q T L A G R T S A I R A M P V E V	440
1381	TCTCAAACATCTACCAGATCTCCCGCAACAGGCGTACCGCCTCAAGTTTAGCCGGTCCG	1440
441	I S N I Y Q I S R E Q A Y R L K F S R S	460
1441	AAACCACCCTCTCCGCCCCGAGAACCAAGGGCGCCAAAGGAGGGAAATGGCTGCTtaag	1500
461	E T T L F R P E N Q G R Q R R E M A A	479
1501	ttgaaaaaacaacagtagtaagcatggcataatgatcacggttatatcatgtaatgt	1560
1561	gaaggagtttaggtgttttttttaaaataattaggatggataacaaaaataaaaggta	1620
1621	gtcctttagaaggtcccttgtttttgaaggtgtttgttgagaggattggagctgttatgt	1680
1681	ataaacttgtcaatgatcaaatgcaaagttccaacatcaaaaaaaaaaaaaaaaaaaaaa	1740
1741	aa	

Figure 2. The nucleotide and predicted amino acid sequence of cDNA clone 8B14. Designations are the same as those listed in Figure 1.

1 aaaattatcagaatntaggtgtaacaaaggtgaagccctcattgccatgcattttgaaaaa 60
61 tggccaaattataaaaattatgtgtcactctttcatctttctcatttttctataaatta 120
121 acctcttcaacacatacattctccattgccacaaattcaatatcctccctccaaaaaa 180
181 acaacaaaaaaagtaaaaaATGGCTAAGTCTACTACTACATTGTTTCTTCTTAG 240
1 M A K S T T T L F L L S 12
241 TTGTTCAATAGCACTAGTGTGTGTAATGGGTGCATGGGCCAAGGAAGGATGCGAGAGAT 300
13 C S I A L V L L N G C M G* Q G R M R E M 32
301 GCAAGGCAATGAGTGCCAAATGACCGGCTTACCGCCCTCGAACCGACGAACCGGATTCA 360
33 Q G N E C S Q I D R L T A L E P T N R I Q 52
361 GGCTGAGGGTGGATTGACTGAGGTGTGGGACACCCAAGACCAGCAGTTCCAATGCTCCGG 420
53 A E G G L T E V W D T Q D Q Q F Q C S S G 72
421 TGTGTCTGTTATTAGAGTACCATTGAGCCTAATGGTCTTCTTTTGCCTTCTTTACTAG 480
73 V S V I R R T I E P N G L L L P S F T S 92
481 TGGCCCTGAGCTCATTACATTGAGCAAGGtaacaaattttaatacaagttctacactaa 540
93 G P E L I Y I E Q 101
541 ttattatttattataatattgagtaatttattatttataagataattacatattcac 600
601 tacacgtgacaattttggcctgataaaaatctttatattacattataaaaaataattatt 660
661 gatatttctactacacgtgacagttttcggcttgataaagcctttatattacattcaaaa 720
721 ataaattattggatgtgaattgtataattattgaaaagtgatgaatatttgtagGGAAT 780
102 G N 103
781 GGAATAAGTGGGCTAATGATCCCGGGATGCCCGGAGACATTGTAATCAATGTCACAAGAA 840
104 G I S G L M I P G C † P E T F E S M S Q E 123
841 TTATGGAGAGAGGGTATGGAGCGAGGAATGAGGGCGGTCGCTTCCAAGACCAACACCAG 900
124 L W R E G M E R G M R G G R F Q D Q H Q 143
901 AAGATCAGGCACCTCCGTC AAGGCCACATTTTCGCAATGCCGGCTGGAGTTGCTCACTGG 960
144 K I R H L R Q G H I F A M P A G V A H W 163
961 GCTTACAACACCGAAATGAGCCTCTTGTGTCTGTATCCTCATTGACACCTCTAACCAT 1020
164 A Y N T G N E P L V A V I L I D T S N H 183
1021 GCTAACCAACTTGACAAGGATTATCCCAAGgttattattatcctcatttctcgtctttt 1080
184 A N Q L D K D Y P K 193
1081 ttaattgatacattatttctaacttaactctttacacattattttttgtcattaagtca 1140
1141 aagtaaaatgtagcacaatatattttacgtgcccacaaatataacgttttttttaagtaat 1200
1201 gtgtatattaataataaccacgtacttctattgtatttaatttaattcattgtattatata 1260
1261 gcagAGATTCTACCTAGCTGGTAAACCCCAACAAGAGCACAGTCCGCCACCAGCACAGAGG 1320
194 R F Y L A G K P Q Q E H S R H Q H R G 212
1321 TGGAGAATCCAGAGGGGTGAACCGCGCAGCGGCCAAGTATTTCAGTGGGCTAGGC 1380
213 G E S Q R G E R G S G G N V F S G L G T 232
1381 CAAGACCATAGCTCAATCCTTCGGAGTTAGTGAGGACATAGCCGAGAAGTCCAAGCCGA 1440
233 K T I A Q S F G V S E D I A E K L Q A E 252
1441 GCAAGCAGAGAGAGGCAACATAGTCTTAGTGCAAGAGGGTCTTCATGTGATCAAGCCCC 1500
253 Q D E R G N I V L V Q E G L H V I K P P 272
1501 AAGCAGCAGTTCATATGATGATGAGAGGGGAACAACGCCCATCGTAGCCAAGGTCTAA 1560
273 S S R S Y D D E R E Q R R H R S P R S N* 292
1561 CGGCTTGGAAAGAGACCATATGCTCGGCTCGTCTTAGTGAGAACATCGACGAACCTCTAA 1620
293 G L E E T I C † S A R L S E N I D E P S K 312
1621 GGCCGATGTTTACTCCCCGAAGCTGGTAGACTTACTACCCTTAACAGCTTCAACCTCCC 1680
313 A D V Y S P E A G R L T T L N S F N L P 332
1681 CATCCTCAGCAACCTCCGCTTGAGTGTGAGAAGGGTGTCTTTACAGGgtatttaatca 1740
333 I L S N L R L S A E K G V L Y R 348
1741 gggctacatataagcattaataaattatttccagacatcctaaaataatattctgatta 1800
1801 atgcaggggttcttactaaatttaatttttgatgcagAACCGGATCATGGCACCACTA 1860
349 N A I M A P H Y 356
1861 CAACCTAAACGCCACAGCATAATCTACGGTGTACGAGGAAGAGGACGTATCCAAATCGT 1920
357 N L N A H S I I Y G V R G R G R I Q I V 376
1921 GAACGCACAAGGAACTCGGTGTTTCGACGACGAGCTAAGACAAGGACAACCTAGTGGTAGT 1980
377 N A Q G N S V F D D E L R Q G Q L V V V 396
1981 CCCACAGAACTTCGCAGTGGTGAAGCAAGCCGGTGAGGAAGGGTTCGAGTGGATCGCCTT 2040
397 P Q N F A V V K Q A G E E G F E W I A F 416
2041 CAAGACATGTGAGAACGCTTGTTCCAAACCTAGCAGGCCGAACATCCGCCATCCGTGC 2100
417 K T C E N A L F Q T L A G R T S A I R A 436
2101 CATGCCCTTAGAAGTGATCTCAAACATCTACCAGATCTCCCGTGAACAGGCGTACCGCCT 2160
437 M P L E V I S N I Y Q I S R E Q A Y R L 456
2161 CAAGTTTAGCCGGTCCGAAACCACCTCTTCCGCCCGGAGAACCAGGGGCCAAAGGAG 2220
457 K F S R S E T T L F R P E N Q G R Q R R 476
2221 GGATTTGGCTGCTtaagttggaaaaaaccaaatgtatgtatgcatggcataatgggtcaca 2280
477 D L A A 480
2281 ttatatatcatgtaattgtgaaggatttttaggtgtttttta 2322

Figure 3. The nucleotide and predicted amino acid sequence of the genomic 11S gene from BAC clone 77L9. The *, § and ‡ indicate the same as those listed in Figure 1. Underlined nucleotides and amino acids indicate differences in the coding regions between sequences 77L9 and 17B7. Introns and untranslated regions are indicated in lowercase.

1	ggtaagccctcattgccatgcattttgaagaatggccaaactatcaaaattatgtgtcac	60
61	tctttcatctttctcatttttctataaattcacctcttcaacacatacattttccattc	120
121	gccacaaattaaatacccttctccaaaaatacaagaaaaaaaaagagataaaaATGGCT	180
1		M A
181	AAGTCTACTACTACATTGTTTCTTCTTAGTGTTCATAGCACTAGTGTGTTGAATGGG	240
3	K S T T T L F L L S C S I A L V L L N G	22
241	TGCATGGGCCAAGGAAGGATGCGAGAAATGCAAGGCAATGAGTGCCTCAATCGACCGCTT	300
23	C M G* Q G R M R E M Q G N E C S Q I D R L	42
301	ACCGCCCTCGAACCGACGCATCGGATTCAGGCTGAGGGTGGGTTGACTGAGGTGTGGGAC	360
43	T A L E P T H R I Q A E G G L T E V W D	62
361	ACCCAGGACCAGCAGTTCCAATGCTCCGGTGTGTCTGTTATTAGCGGTACCATTGAGCCT	420
63	T Q D Q Q F Q C S S G V S V I R R T I E P	82
421	AATGGTCTTCTTTGCCTTCTTCTACTAGTGGCCCTGAGCTCATTTACATTGAGCAAGgt	480
83	N G L L L P S F T S G P E L I Y I E Q	101
481	aacaaatattaatacaagttctacaataatcattataataatacggagtaatttataat	540
541	ttatattataaggtaattaattacgtattgagtaacttgtgagatatttcggctcaaaaat	600
601	aacattcatatgttatattgtgtagattattgaaaagtgatagaattttttagtagGGAATG	660
102		G N
661	GAATAAGTGGGCTGATGATCCCGGGTGCCTCCGAGACATTTGAATCAATGTCACAAGAAT	720
104	G I S G L M I P G C † P E T F E S M S Q E	123
721	CATGGAGAGAGGGTATGGAGCGAGGAATGAGGGGCGGTGCTTCCAAGACCAACACCAGA	780
124	S W R E G M E R G M R G G R F Q D Q H Q	143
781	AGATCAGGCACCTCCGTCAAGGCCACATTTTCGCAATGCCGGCCGAGTTGCTCACTGGG	840
144	K I R H L R Q G H I F A M P A G V A H W	163
841	CTTACAACCTCCGAAATGAGCCACTTGTGTCTGTTATCCTCATTGACACCTCTAACCATG	900
164	A Y N S G N E P L V A V I L I D T S N H	183
901	CTAACCAACTTGACAAGGATTACCCCAAGgtacgttacacttcaataatattccttatta	960
184	A N Q L D K D Y P K	193
961	cctcgtctttttgttacatatttcttattctttagtctttttattcgtttacaaatttaca	1020
1021	attattcttggcaagcgcttctccaatatttagactcaataatgtgtcaaaatataatg	1080
1081	tagcaacaataaaaagactagagaaaataatgtaacggttttttaaaatgatgcttacgta	1140
1141	ccaacttcaacaataattaataacaccacttgtgcatttgatcaatgatttatatgacagA	1200
1201	GATTCTACCTAGCTGGTAAACCCCAACAAGAGCACAGTCCGCCACCATCACAGGGTGGAG	1260
194	R F Y L A G K P Q Q E H S R H H H R G G	213
1261	AATCCCAGAGGGGTGAACACGGCAGCGACGGCAACGTATTCAGTGGCCTAGACACCAAGT	1320
214	E S Q R G E H G S D G N V F S G L D T K	233
1321	CCGTAGCTCAATCCTTCGGAGTTAGTGAGGACATAGCGGAGAAGCTCCAAGCCAAGCAGG	1380
234	S V A Q S F G V S E D I A E K L Q A K Q	253
1381	ACGAGAGAGGCAACATAGTCTTAGTGCAAGAGGGTCTTCATGTGATCAAGCCCCAAGCA	1440
254	D E R G N I V L V Q E G L H V I K P P S	273
1441	GCAGGTCATATGATGATGAGAGGGAACAGCGCCATCGTAGCCCAAGGTCTAACGGCTTGG	1500
274	S R S Y D D E R E Q R H R S P R S N* G L	293
1501	AAGAGACCATATGCTCGGCTCGCCTTAGTGAGAACATCGACGATCCCTCTAAGGCCGATG	1560
294	E E T I C † S A R L S E N I D D P S K A D	313
1561	TTTACTCCCCGGAAGCTGGTAGACTTACCACCCTTAACAGCTTCAACCTCCCCATCCTCA	1620
314	V Y S P E A G R L T T L N S F N L P I L	333
1621	GCAACCTCCGCTTAGTGCTGAGAAGGGTGTCTCTACAGGgtaattaatcagggtata	1680
334	S N L R L S A E K G V L Y R	347
1681	tataagtattaaatatatttttccgtatatcctaaaatagtaatctgtcagtagaact	1740
1741	ttaactaaattatttttgatgcagAACGCAATCATGGCACCACTACAACCTAAACGCC	1800
348		N A I M A P H Y N L N A
1801	CACAGCATAATCTATGGTGTACGAGGACGAGGACGTATCCAATCGTGAACGCACAAGGA	1860
360	H S I I Y G V R G R I Q I V N A Q G	379
1861	AACTCAGTGTTTGACGATGAGCTAAGACAAGGACAAGTGTGGTAGTCCCACAGAAGTTC	1920
380	N S V F D D E L R Q G Q L V V V P Q N F	399
1921	GCAGTGGTGAAGCAAGCCGGTGAAGGAGGATTCGAGTGGATCGCCTTCAAGACATGTGAG	1980
400	A V V K Q A G E E G F E W I A F K T C E	419
1981	AACGCTTTGTTCCAAACCCTAGCAGGCCGAACATCCGCCATACGTGCCATGCCGTAGAA	2040
420	N A L F Q T L A G R T S A I R A M P V E	439
2041	GTAATCTCAAACATCTACCAGATCTCCCGCGAAGCAGGCGTACCGCCTCAAGTTTAGCCGG	2100
440	V I S N I Y Q I S R E Q A Y R L K F S R	459
2101	TCCGAAACCACCTCTTCCGCCCCGAGAACCAAGGGCGCCAAAGGGGAAATGGCTGCT	2160
460	S E T T L F R P E N Q G R Q R R E M A A	479
2161	taagttgaaaaacaacaacgtagtaagcattggcataatgatcacgttatatatcatgta	2220
2221	atgtgaaggagtttttaggttgttttttttaaaataattaggatggataacaaaataaaa	2280
2281	ggtagtcctttagaaggtcccttgtttttgaaggtgtttttgttgagaggattggagctgtt	2340
2341	atgta	

Figure 4. The nucleotide and predicted amino acid sequence of the genomic 11S gene from BAC clone 164F2. Designations are the same as those listed in Figure 3. Underlined nucleotides and amino acids indicate differences between the 164F2 and 8B14 sequences.

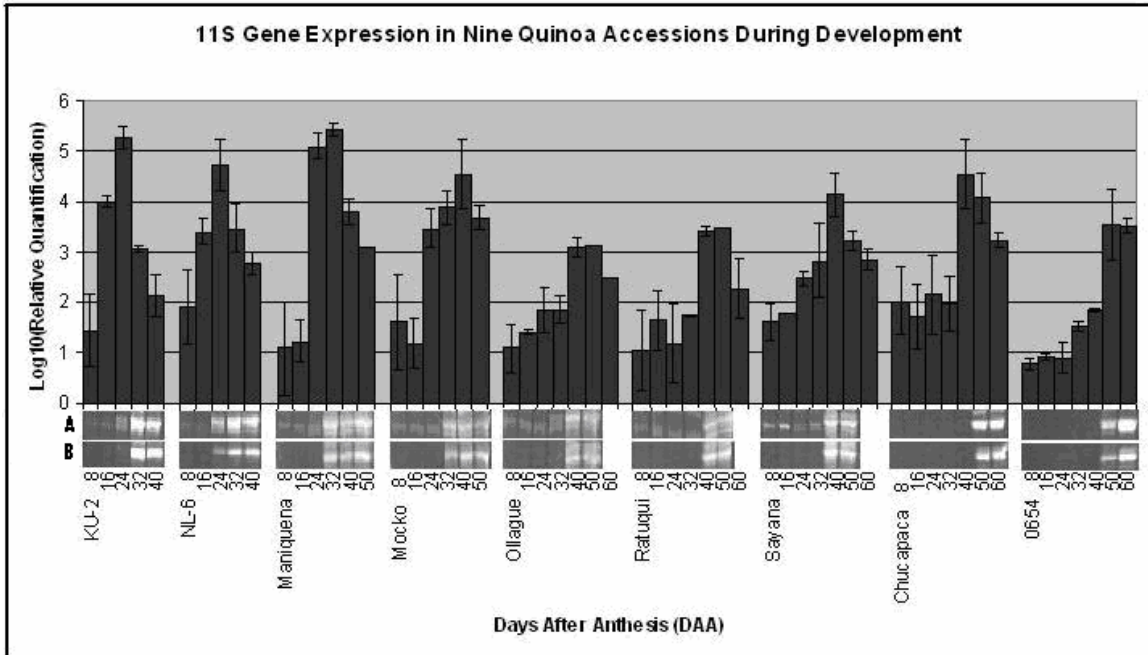


Figure 5. 11S gene expression and protein accumulation during the development of quinoa seeds. Quinoa accessions and days after anthesis (DAA) are listed on the horizontal axis. Gene expression was measured in $\text{Log}_{10}(\text{RQ})$ units using the GAPDH gene as an endogenous control. The results of the SDS-PAGE analysis are shown below the graph. A represents the acidic subunit of the 11S polypeptide and B represents the basic subunit of the 11S polypeptide.

Species	Position	Sequence	Position
Quinoa *8B14	291	NGLEETICSAHLSENIDDPDK-ADVYSPEAGRLTTLNSFNLPIILSNLRLAEKGVLY	347
Quinoa *17B7	292	NGLEETICSAHLSENIDDPDK-ADVYSPEAGRLTTLNSFNLPIILSNLRLAEKGVLY	348
Amaranth	298	NGVEETICSAHLAVVDDPSK-ADVYTPPEAGRLTTLNSFNLPIILRHLRLAAKGVLY	354
Hazelnut	320	NGFEEITCSLRLRENICTRSP-ADIYTEVGRINLVNSHLPVLRWLQLAEKGVLY	376
English Walnut	315	NGLEETICTLRLRENIGDPSK-ADIYTEEAGRISTVNSHLPVLRWLQLAEKGVLY	371
Magnolia	288	NGLEETQCSAKLTYIADDPK-ADVYNPQAGRITSLNSQFFILVLRQLAEKGVLY	344
Cashew	271	NGIEETICTMRLKENINDPDK-ADIYTPPEVGRITTLNSLNLPIILRWLQLAEKGVLY	327
English Oak	298	NGIEETLCTLRLENIHDPDK-ADIYNPQAGRISTLNSHNLVLRWLQLAEKGVLY	354
Sesame	281	NGLEETLCTLRLENLDEPK-ADVYNPHGGRISLNSLNLVLRWLQLAEKGVLY	337
Brazil Nut	279	NGLEETICSAFIQNIIDPDK-ADVYNPQAGRITTLVNSLNVPIILVLRQLAEKGVLY	335
Castor Bean	289	NGVEETICTMRLKENIADDPK-ADVYVPEVGRVSTVNSHNLPIILRWLQLAEKGVLY	345
Japanese Chestnut	353	NGIEETLCTLRLENIHDPDK-ADIYNPQAGRISTLNSHNLVLRWLQLAEKGVLY	409
Cotton	324	NGLEETFCSMRIKENIADDPK-ADIFNPQAGRISTLNRFLVLRWLQLAEKGVLY	380
Almond	314	NGLEETFCSLRLKENIGDPSK-ADIFSPRAGRISTLNSHNLVLRWLQLAEKGVLY	370
Pumpkin	296	NGLEETICTLRLKQNIIDPDK-ADVFNPRGGRISTANYHLPVLRWLQLAEKGVLY	352
Coffee	304	NGLEETLCTVRLSENIGLQPK-ADVFNPRAGRITVNSQIPVLRWLQLAEKGVLY	360
Radish	289	NGLEETICSMRTHENIDDPK-ADVYKPNLGRVTVNSYLPVLRWLQLAEKGVLY	345
Rape	299	NGLEETICSAKCTDNLDPPK-ADVYKQPLGYISTLNSYDLPVLRWLQLAEKGVLY	355
Fava Bean *B4	303	NGLEETICSLKIRENIADDPK-ADLYNPRAGSISTANSLVLRWLQLAEKGVLY	359
Fava Bean *A2	315	NGLEETVCTAKLRLNIGSSSS-EDIYNPQAGRIKTVPSLDLPVLRWLQLAEKGVLY	371
Tare *LegB	304	NGLEETICSAKIRENIADDPK-ADLYNPRAGRISTANSLVLRWLQLAEKGVLY	360
Tare *LegA	313	NGLEETVCTAKLRLNIGSSSS-EDIYNPQAGRIKTVPSLDLPVLRWLQLAEKGVLY	369
Norbonne Vetch	297	NGLEETVCTAKLRLNIGSSSS-EDIYNPQAGRINTVPSLDLPVLRWLQLAEKGVLY	353
Rice	302	NGLEENFCTIIVRVNIENPDK-ADSYNPRAGRITVNSQKFFILNLIOMAEKGVLY	358
Thale Cress	282	NGLEETICSAKCTDNLDPPK-ADVYKQPLGYISTLNSYDLPVLRWLQLAEKGVLY	338
Wild Soybean	344	NGVEENICTMRLKENIADDPK-ADLYNPRAGRISTLNSLNLVLRWLQLAEKGVLY	400
Soybean *A2B1a	300	NGIDEETICTMRLKRONIGSSSS-EDIYNPQAGSITVPSLDLPVLRWLQLAEKGVLY	356
Soybean *A5A4B3	378	NGVEENICTMRLKENIADDPK-ADLYNPRAGRISTLNSLNLVLRWLQLAEKGVLY	434
White Mustard	320	NGLEETICSMRTHENIDDPK-ADIYKPNLGRVTVNSYLPVLRWLQLAEKGVLY	376
Sunflower	305	NGVEETICSMRFKVIIDNPSK-ADVFNPAAGSIALNSFVPIILRHLRLAEKGVLY	361
Navel Orange	299	NGFEEITICTMRLRHNIDDPK-ADVYNPQAGRITVNSHNLVLRWLQLAEKGVLY	355
Buckwheat	313	NGLEAFFCNLKFRONVNRPSK-ADVFNPRAGRINTVDSNLPVLRWLQLAEKGVLY	369
Oat	317	NGLEENFCSLEARONIENPDK-ADLYNPRAGRITLHNSKLPVLRWLQLAEKGVLY	373
African Oil Palm	286	NGLEVAMCSMRNENIDSSRR-ADVYIPRGGRIITLNSQRLPMLVLRWLQLAEKGVLY	342
Wild Ginger	278	NGMEETICTMRLKRONIGSSSS-EDIYNPQAGRINTVPSLDLPVLRWLQLAEKGVLY	334
Pea *LegA	332	NGLEETVCTAKLRLNIGSSSS-EDIYNPQAGRIKTVPSLDLPVLRWLQLAEKGVLY	388
Pea *LegK	169	NGLEETICSAKIRENIADDPK-ADLYNPRAGRISTANSLVLRWLQLAEKGVLY	225
Arrowhead	392	NGIEETICNLKFKVIIGNPIH-ADVYSREGGHLLTLNSFNLPIILSYLQLAEKGVLY	448
Ginkgo	273	NNVEEFYCSMRRLRHNIDDPK-ADVYVRRGGRLNTVNSLNLVLRWLQLAEKGVLY	329
Beefsteak Mint	285	NGLEEFCSMRKIMSNDNPRE-ADVYSRQAGKLVVDMHRLVLRWLQLAEKGVLY	341
Lupine	327	NGLEETLCTMRLRHNIGESTS-EDAYNPQAGRIFKTLTSDIFILGLGLAAEHGVIY	383
Peanut	298	NGIEETICTASVKKNIIGNRS-EDIYNPQAGSLKLTANDLNLVLRWLQLAEKGVLY	354
Chickpea	311	NGFEEICTALRHQNIIDPPK-ADVYNPQAGRIKTVPSLDLPVLRWLQLAEKGVLY	367
Douglas Fir	315	NDVEEVVVCALRVKHNADNPED-ADIYVRDGGRLNIVNRFVLRWLQLAEKGVLY	371
Dawn Redwood	316	GGLHGFYCNMRLRHNADRPDD-ADIFVRDGGRLNIVNRFVLRWLQLAEKGVLY	372
Japanese Cedar	325	NGLVVLFNCMRLRHNADNPED-ADVYVRDGGRLNIVNRFVLRWLQLAEKGVLY	381
Yam	293	NGLEEAICYAVVQYVLDPEEDSDVYSRQAGRLKQVDLNLVLRWLQLAEKGVLY	350
White Spruce	311	NGVEELVCPVLRVKNADNPED-ADVYVRDGGRLNIVNRFVLRWLQLAEKGVLY	367
Corn	290	DVDRHNVCAMEVVRSVERLDG-ADVYSPGAGRITLTSHPFVLRWLQLAEKGVLY	346
Incense Cedar	314	GGLHGFYCNMRLRHNADRPDD-ADIFVRDGGRLNIVNRFVLRWLQLAEKGVLY	370
Eastern White Pine	287	NGVEELVCPVLRVKNADNPED-ADVYVRDGGRLNIVNRYVLRWLQLAEKGVLY	343
Welwitschia	321	GVAEETVCSMRMRHFDNPNP-ADVYVAGGGRMIVNRYVLRWLQLAEKGVLY	377
Ephedra	319	GFESEVVCNQRIRHNINRQD-EDFYHPRAGFMSVANSFLVLRWLQLAEKGVLY	375
Buko	368	GVAEEGSCSMRLRQSLNRPD-ADIYVRGAGRVNLANALVLRWLQLAEKGVLY	424
Wheat	380	NGLEENFCDHKLIVNIDDPK-ADIYNPRAGRITLNSQFFILNLIOMAEKGVLY	436

Figure 6. Alignment using ClustalW of 55 legumin amino acid sequences from 50 different species. The amino acid alignment was used as a template for the CDS alignment. Only approximately 56 amino acid residues are shown which make up the N-terminal sequence of the basic subunit and in most cases, the C-terminal-asparagine residue of the acidic subunit. The * indicates amino acid residues that are the same in all sequences. The numbers on either side of the alignment indicate amino acid position.

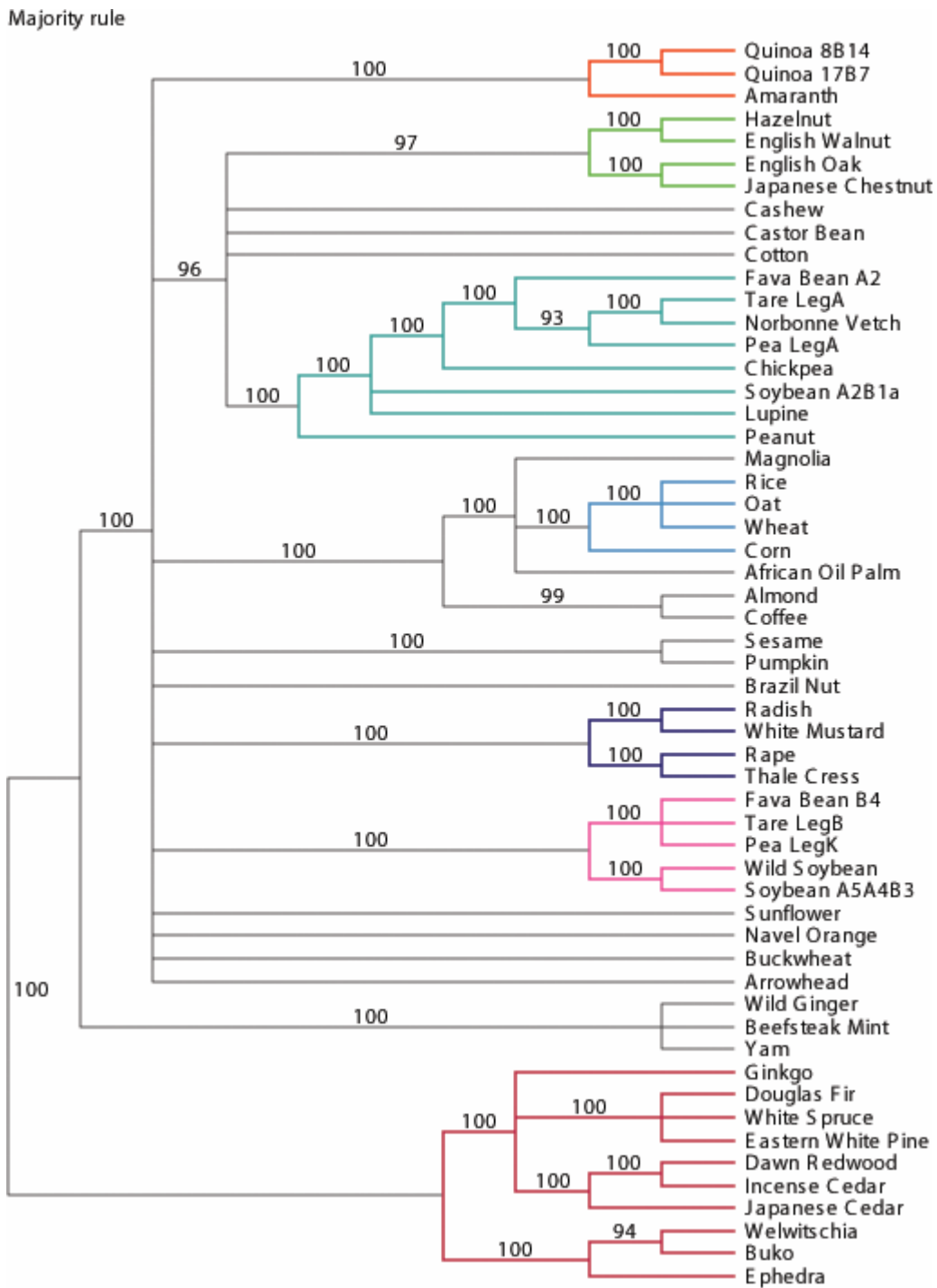


Figure 8. Bayesian consensus tree reconstruction of coding DNA sequences of the legumin basic subunit. Unresolved portions of the tree represent posterior probabilities below 90. Colored portions represent monophyletic groups identified by both Bayesian and Parsimony tree reconstructions.

Table 1. Amino acid composition of the two loci of the 11S gene in quinoa 8B14 and 17B4 compared to analogous genes in rice, corn and soybean

Amino Acid Composition					
Amino acid	<i>C. quinoa</i> 8B14	<i>C. quinoa</i> 17B7	Amino acid	<i>C. quinoa</i> 8B14	<i>C. quinoa</i> 17B7
Ala	6.39	6.59	Leu*	7.27	7.69
Arg	8.59	9.01	Lys*	2.86	2.86
Asn	5.07	5.27	Met*	2.20	1.98
Asp	3.96	3.52	Phe*	3.52	3.52
Cys*	1.10	1.10	Pro	4.41	4.40
Glu	8.37	8.35	Ser	7.93	7.47
Gln	7.49	7.69	Thr*	3.74	4.18
Gly	8.81	9.23	Trp*	0.88	0.88
His*	3.08	2.64	Tyr*	2.64	2.42
Ile*	6.17	6.37	Val	5.51	4.84

*Essential amino acids

Table 2. Quinoa accession, location and maturation type for which 11S gene and protein expression data were obtained

Accession	Location*	Maturation Type
KU-2	Coastal	Type I
NL-6	Coastal	Type I
Maniqueña	Salares	Type II
Mocko	Salares	Type II
Ollague	Salares	Type III
Ratuqui	Altiplano	Type III
Sayaña	Altiplano	Type III
Chucapaca	Altiplano	Type IV
0654	Valley	Type V

*Valley and Salares are subdivisions of the Altiplano type

Table 3. Organisms' common name, species, family, order and GenBank accession number from which the 11S gene cDNA sequences were obtained

Common Name	Species	Family	Order	Accession Number
Arrowhead	<i>Sagittaria sagittifolia</i>	Alismataceae	Alismatales	Y09116 .
African Oil Palm	<i>Elaeis guineensis</i>	Arecaceae	Arecales	AF261691 .
Sunflower	<i>Helianthus annuus</i>	Asteraceae	Asterales	M28832 .
Thale Cress	<i>Arabidopsis thaliana</i>	Brassicaceae	Brassicales	NM_123779 .
Rape	<i>Brassica napus</i>	Brassicaceae	Brassicales	M16860 .
Radish	<i>Raphanus sativus</i>	Brassicaceae	Brassicales	X59808 .
White Mustard	<i>Sinapsis alba</i>	Brassicaceae	Brassicales	AY846388 .
Quinoa *17B7	<i>Chenopodium quinoa</i>	Amaranthaceae	Caryophyllales	AY562549 .
Quinoa *8B14	<i>Chenopodium quinoa</i>	Amaranthaceae	Caryophyllales	AY562550 .
Amaranth	<i>Amaranthus hypochondriacus</i>	Amaranthaceae	Caryophyllales	X82121 .
Buckwheat	<i>Fagopyrum esculentum</i>	Polygonaceae	Caryophyllales	AF152003 .
Incense Cedar	<i>Calocedrus decurrens</i>	Cupressaceae	Coniferales	X95540 .
Japanese Cedar	<i>Cryptomeria japonica</i>	Cupressaceae	Coniferales	X95542 .
Dawn Redwood	<i>Metasequoia glyptostroboides</i>	Cupressaceae	Coniferales	X95544 .
White Spruce	<i>Picea glauca</i>	Pinaceae	Coniferales	X63192 .
Eastern White Pine	<i>Pinus strobus</i>	Pinaceae	Coniferales	Z11486 .
Douglas Fir	<i>Pseudotsuga menziesii</i>	Pinaceae	Coniferales	L07484 .
Pumpkin	<i>Cucurbita pepo</i>	Cucurbitaceae	Cucurbitales	M36407 .
Yam	<i>Dioscorea caucasica</i>	Dioscoreaceae	Dioscoreales	X95510 .
Ephedra	<i>Ephedra gerardiana</i>	Ephedraceae	Ephedrales	Z50777 .
Brazil Nut	<i>Bertholletia excelsa</i>	Lecythidaceae	Ericales	AY221641 .
Peanut	<i>Arachis hypogaea</i>	Fabaceae	Fabales	AF125192 .
Chickpea	<i>Cicer arietinum</i>	Fabaceae	Fabales	Y15527 .
Soybean *A2B1a	<i>Glycine max</i>	Fabaceae	Fabales	D00216 .
Soybean *A5A4B3	<i>Glycine max</i>	Fabaceae	Fabales	AB195712 .
Wild Soybean	<i>Glycine soja</i>	Fabaceae	Fabales	X79467 .
Lupine	<i>Lupinus albus</i>	Fabaceae	Fabales	AJ938034 .
Pea *LegA	<i>Pisum sativum</i>	Fabaceae	Fabales	AJ132614 .
Pea *LegK	<i>Pisum sativum</i>	Fabaceae	Fabales	X07015 .
Fava Bean *A2	<i>Vicia faba</i>	Fabaceae	Fabales	X55014 .
Fava Bean *B4	<i>Vicia faba</i>	Fabaceae	Fabales	X14237 .
Norbonne Vetch	<i>Vicia narbonensis</i>	Fabaceae	Fabales	Z46803 .
Tare *LegA	<i>Vicia sativa</i>	Fabaceae	Fabales	Z32835 .
Tare *LegB	<i>Vicia sativa</i>	Fabaceae	Fabales	Z32796 .
Hazelnut	<i>Corylus avellana</i>	Betulaceae	Fagales	AF449424 .
Japanese Chestnut	<i>Castanea crenata</i>	Fagaceae	Fagales	AF525749 .
English Oak	<i>Quercus robur</i>	Fagaceae	Fagales	X99539 .
English Walnut	<i>Juglans regia</i>	Juglandaceae	Fagales	AY692446 .
Coffee	<i>Coffea arabica</i>	Rubiaceae	Gentianales	AF054895 .
Ginkgo	<i>Ginkgo biloba</i>	Ginkgoaceae	Ginkgoales	Z50778 .
Buko	<i>Gnetum gnemon</i>	Gnetaceae	Gnetales	Z50779 .
Beefsteak Mint	<i>Perilla frutescens</i>	Lamiaceae	Lamiales	AF180392 .
Sesame	<i>Sesamum indicum</i>	Pedaliaceae	Lamiales	AF240004 .
Magnolia	<i>Magnolia salicifolia</i>	Magnoliaceae	Magnoliales	X82464 .

Castor Bean	<i>Ricinus communis</i>	Euphorbiaceae	Malpighiales	AF262998.
Cotton	<i>Gossypium hirsutum</i>	Malvaceae	Malvales	M16905.
Wild Ginger	<i>Asarum europaeum</i>	Aristolochiaceae	Piperales	X95508.
Oat	<i>Avena sativa</i>	Poaceae	Poales	X17637.
Rice	<i>Oryza sativa</i>	Poaceae	Poales	XM 464834.
Wheat	<i>Triticum aestivum</i>	Poaceae	Poales	S62630.
Corn	<i>Zea mays</i>	Poaceae	Poales	AF371279.
Almond	<i>Prunus dulcis</i>	Rosaceae	Rosales	X78119.
Cashew	<i>Anacardium occidentale</i>	Anacardiaceae	Sapindales	AF453947.
Navel Orange	<i>Citrus sinensis</i>	Rutaceae	Sapindales	U38914.
Welwitschia	<i>Welwitschia mirabilis</i>	Welwitschiaceae	Welwitschiales	Z50780.

* Indicates the type of subunit

Appendix

Relative quantification of mRNA using one-step real time RT-PCR

Quantitative real-time PCR is a technique capable of recording the progress of the PCR throughout the process and therefore determine quantities of specified starting materials. Reverse transcriptase (RT) real-time PCR is a method used to quantify RNA transcripts and so is useful in gene expression assays. Relative quantification (RQ) determines the change in expression of a target gene sequence in reference to an appropriate endogenous control gene. RQ is capable of determining the initial quantities of specific transcripts without knowing the total initial RNA concentration and without the use of a standard curve that is necessary for absolute quantification.

There are several components that are necessary for an RQ experiment. The target is the sequence under study. An endogenous control is a gene with a constant level of expression in order to normalize the quantification of the target. Housekeeping genes such as glyceraldehyde-3-phosphate dehydrogenase (GAPDH), ribosomal RNA (rRNA) and β -actin are commonly used as endogenous controls due to their stable expression levels. Each sample must include a target and an endogenous control. A calibrator is a sample which serves as a basis for comparison for all the other samples within the experiment. In addition, sample replication is necessary to ensure statistical significance.

General Outline:

- A. RNA Preparation
- B. Primers and Probe Selection
- C. Multiplexing vs. Singleplexing
- D. RQ RTPCR cocktail for multiplexing
- E. RQ RTPCR cocktail for singleplexing
- F. Creating a RQ Plate Document using Sequence Detection Systems Software
- G. Thermocycler conditions
- H. Creating an RQ Study

A. RNA Preparation

1. Collect plant material and immediately freeze in liquid nitrogen.
2. Store plant material at -80°C.
3. Extract RNA using a RNeasy® Plant Mini Kit (Qiagen, Valencia, CA) or appropriate protocol and follow the manufacturer's instructions.
4. Quantify total RNA.
5. Adjust RNA concentration to 50 µg/ml using RNase free water.

Note: The total RNA concentration may need to be increased or decreased based on the quantity of target mRNA present in the total RNA.

B. Primers and Probe Selection

There are several different methods for selecting primers and probes for the target sequence and the endogenous control. Assays-by-DesignSM is a service provided by Applied Biosystems (Foster City, CA) in which only a sequence is provided and primer and probe sets are designed, synthesized, and delivered in the appropriate concentrations. Primer Express® Software is designed to assist in selecting primers and probes from a given sequence. Primer and probe sets are necessary for the target and endogenous control sequences.

Note: Multiplexing requires the probe for the target sequence and the probe for the endogenous control to have two different dyes. In singleplexing experiments, the probe for the target sequence and the probe for the endogenous control may have the same dye. In multiplexing, FAM and VIC dyes are commonly used. In singleplexing, the FAM dye may be used for both

C. Multiplexing vs. Singleplexing

A PCR experiment can either be conducted as a singleplex in which only a single primer set is present or a multiplex reaction in which two or more primer sets are present within a sample. In a singleplex experiment, the primers and probes specific to the target sequence and those for the endogenous control are in separate wells or tubes while multiple primer sets for single or multiple targets and endogenous controls may be in a single well or tube in a multiplex experiment. Multiplexing requires less sample, less reagents and minimizing pipetting error. However, factors such as the number of targets or the abundance of the endogenous control must be taken into consideration. As the

number of targets increase, the multiplexing method becomes less effective. In addition, the endogenous control must be more abundant than the targets.

Materials:

RNA samples

96-well optical reaction plate

Optical adhesive cover

TaqMan® Universal PCR Master Mix with ROX (2X)

20X Assay Mix for target sequence

Forward Primer Concentration (μM): 18 μM

Reverse Primer Concentration (μM): 18 μM

Probe Concentration (μM): 5 μM

20X Assay Mix for endogenous control

Forward Primer Concentration (μM): 18 μM

Reverse Primer Concentration (μM): 18 μM

Probe Concentration (μM): 5 μM

Multiscribe™ Reverse Transcriptase

RNase-free water

Instructions

1. In a 96-well optical reaction plate, apply 10 μl RNA sample (50 $\mu\text{g}/\text{ml}$) to each well in duplicates or triplicates for statistical significance.
2. Prepare cocktail sufficient for the number of wells to be used on ice.

3. Add 15µl of cocktail (see instructions below) to each well.
4. Cover with an optical adhesive cover and ensure wells are sealed to avoid evaporation during the PCR.
5. Centrifuge briefly to ensure sample is at the bottom of each well.
6. Keep plate on ice until ready for use.

RQ RT-PCR cocktail for multiplexing

One Well Cocktail	Volume (µl)
1. TaqMan® Universal PCR Master Mix with ROX (2X)	12.500
2. 20X Assay Mix for target sequence	0.625
3. 20X Assay Mix for endogenous control	0.625
4. Multiscribe™ Reverse Transcriptase (on ice)	0.500
5. RNase-free water	<u>0.750</u>
Total	15.000

RQ RT-PCR cocktail for singleplexing

Note: Singleplexing experiments require two cocktails to be prepared, one containing the assay mix for the target sequence and the other containing the assay mix for the endogenous control

One Well Cocktail	Volume (µl)
1. TaqMan® Universal PCR Master Mix with ROX (2X)	12.50
2. 20X Assay Mix for target sequence or endogenous control	1.25
3. Multiscribe™ Reverse Transcriptase (on ice)	0.50
4. RNase-free water	<u>0.75</u>
Total	15.00

D. Creating a RQ Plate Document using Sequence Detection Systems Software

These steps are also illustrated in the Relative Quantification Getting Started Guide provided by Applied Biosystems. Refer to the guide for further details.

1. Create a new RQ Plate document:
 - a. Assay: Relative Quantification (ddCt) Plate
 - b. Container: 96-Well Clear
 - c. Template: Bland Document
 - d. Plate Name: Enter plate name
2. Select detectors: Add any dyes that have been used to label probes that will be used in the experiment.
 - a. Multiplexing: generally the probe for the target is labeled with FAM dye and the probe for the endogenous control is labeled with VIC® dye
 - b. Singleplexing: generally only FAM dye is used to label the probes of both the target and endogenous control.
3. Determine the detectors and tasks of each well.
 - a. The task will be either to detect target or endogenous control or both (if multiplexing).
4. Enter sample names using the well inspector (detectors and tasks can also be determined at this step).

E. Thermocycler Conditions for One-Step RT-PCR

1. Stage 1 (1 Rep): 48.0°C for 30 minutes

Note: This stage may be lengthened or shortened depending on the type of reverse transcriptase being used.

2. Stage 2 (1 Rep): 95.0°C for 10 minutes

3. Stage 3 (40 Reps): 95.0°C for 15 seconds

60.0°C for 1 minute

Note: More reps may be used, especially if the quantity of target mRNA is small

4. Sample volume: 25µl

F. Creating an RQ Study

1. Create a new RQ Study document:

- a. Assay: Relative Quantification (ddCt) Study
- b. Container: 96-Well Clear
- c. Plate Name: Enter plate name

2. Add plates to the study.

Note: Only plates that have been run under the same conditions will be allowed to be compared in an RQ Study. Also, samples with the same name will be averaged and shown under a single sample name.

3. RQ Study Document:

- a. RQ Detector Grid: shows detectors used in the experiment.
- b. RQ Sample Grid: shows samples associated with selected detectors.

- i. Sample summary: includes information about detectors, tasks, and RQ calculations.
- ii. Well information: includes information about the location of samples on the plate.
- c. RQ Results Panel: contains information about the plates included in the study, the amplification plot, and relative gene expression.

4. Analysis Settings

- a. Detector: All
- b. Auto Ct
- c. Determine which sample will be used as the calibrator.

Note: A calibrator is a sample which serves as a basis for comparison for all the other samples within the experiment.

- d. Select the endogenous control detector.
- e. Control type: only an option if plates contain multiplexed and singleplexed reactions.
- f. RQ Min/Max Confidence (for error bar calculations): 95.00%
- g. Select 'Remove Outlier'
- h. Click 'OK & Reanalyze'

5. Verify the accuracy of the baseline and threshold.

- a. The amplification curve should include a:
 - i. Plateau phase
 - ii. Linear phase
 - iii. Geometric phase

iv. Background

v. Baseline

- b. The maximum baseline should be set before the amplification curve.
- c. The threshold should be set in the geometric phase of the amplification curve.
- d. Adjust the baseline and threshold manually if necessary by selecting Manual Ct under the Analysis Settings.
- e. Always reanalyze after making any changes in the analysis settings.

6. Export the RQ Study Data

- a. Under file, select export, then results, then sample summary.
- b. Data can be further analyzed in Microsoft® Office Excel.